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(54) Title: COMPOSITION AND METHOD FOR TREATING CANCERS CHARACTERIZED BY OVER-EXPRESSION OF THE C-FMS PROTO-ONCOGENE

(57) Abstract

A composition and method for treating cancers characterized by over-expression of the c-fms proto-oncogene/M-CSF receptor protein are provided. The composition involves an M-CSF polypeptide cross-linked to a cytotoxic agent capable of crossing into the cytoplasm of the cell bearing the receptor and killing the cell.

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COMPOSITION AND METHOD FOR TREATING CANCERS
CHARACTERIZED BY OVER-EXPRESSION OF THE C-FMS PROTO-ONCOGENE

The present invention refers generally to the treatment of a variety of cancers characterized by the over-expression of the protein receptor, c-fms. More specifically, the invention refers to a composition for such treatment including the M-CSF polypeptide linked to a cytotoxic agent.

BACKGROUND OF THE INVENTION

A variety of oncogenes have been associated with specific cancers. The oncogene fms has come under recent scrutiny ~~as being related to breast, lung, pancreatic,~~ ovarian, renal, and possibly other carcinomas, including acute myelocytic leukemia (AML). See, e.g., D. J. Slamon et al, Science, 224:256-262 (1984); C. Walker et al, Proc. Natl. Acad. Sci., USA, :1804-1808 (April 1987). See also, J. H. Ohyashiki et al, Cancer Genet. Cytogenet., 25:341-350 (1987); H. D. Preisler et al, Cancer Research, 47:874-880 (Feb. 1987); C. W. Rettenmier et al, J. Cell. Biochem., 33:109-115 (1987); and R. Sacca et al, Proc. Natl. Acad. Sci. USA, 82:3331-3335 (1986). The product of the c-fms proto-oncogene is believed to be related to, and possibly identical with, a receptor of macrophage colony-stimulating factor (M-CSF). See, e.g., C. J. Sherr et al, Cell, 41:665-676 (1985);

There remains a need in the treatment of such cancers for therapeutic products capable of destroying the carcinoma cells without severely adversely affecting the patient otherwise.

BRIEF DESCRIPTION OF THE INVENTION

As one aspect of the invention there is provided a composition for treating cancers which are characterized by high level expression of the c-fms proto-oncogene/M-CSF

receptor gene. The composition includes M-CSF polypeptide (or the active fragment thereof) crosslinked to a cytotoxic agent, which is capable of crossing the membrane of the cell bearing the c-fms gene product/M-CSF receptor and acting in the cytoplasm to destroy the cell. Preferred cytotoxic agents include A and B chain toxins, A chain toxins and genetically engineered toxins.

In a further embodiment the composition may comprise a monoclonal antibody (or a portion thereof) to c-fms gene product/M-CSF receptor conjugated to a cytotoxic agent. This monoclonal moiety recognizes and binds to the c-fms gene product/M-CSF receptor. Antibody conjugates for the delivery of compounds to target sites and methods for preparing the same are known in the art. See e.g. U.S. Patent 4,671,958.

Still a further aspect of the invention involves a method for making the M-CSF/cytotoxic agent composition. The M-CSF and toxin may be linked by employing one or more heterofunctional or bifunctional protein cross linkers or by genetic fusion. The bifunctional cross-linkers are chosen to ensure that the M-CSF/toxin composition is stable while the composition is homing to the target cell. At the same time the crosslinker has to permit the release of the toxin portion after the M-CSF/toxin composition has entered the cell. See, e.g. Molecular Action of Toxins and Viruses, P. Cohen and S. van Heyningen, eds., Elsevier, New York, pp51-105 (1982).

As another aspect there is disclosed a method for treating cancers characterized by an over-expression of the c-fms proto-oncogene/M-CSF receptor gene. This method involves regionally administering to the in vivo site of such a cancer, the composition of the invention, or, alternatively, administering the composition in an ex vivo purging treatment of a mixture of cells. The composition acts by attaching to the c-fms protein on the carcinoma and delivering the toxin through the cell membrane, where the

toxin destroys the cell. Among such receptor over-expressing cancers are acute myelocytic leukemia, ovarian carcinoma, lung carcinoma, and those recited above.

Other aspects and advantages of the present invention will be apparent upon consideration of the following detailed description of the invention, including illustrative examples of the practice thereof.

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 illustrates a DNA and amino acid sequence for an M-CSF polypeptide.

DETAILED DESCRIPTION OF THE INVENTION

The therapeutic composition of the invention is a conjugate of M-CSF which is capable of binding to the c-fms proto-oncogene/M-CSF receptor gene product on certain cancer cells, and a cytotoxic agent capable of being transported through the cell membrane and acting in the cytoplasm to destroy the cell.

The M-CSF for use in the present invention may be recovered from natural sources and purified. (See e.g., UK Patent 2,016,477 and PCT published application WO86/04587). Alternatively, the M-CSF may be produced recombinantly. One possible recombinant M-CSF polypeptide useful in the present invention has been described in PCT published application WO86/04607. Another M-CSF polypeptide is described in co-pending, co-owned US patent application SN940,362 and in G. G. Wong et al, Science, 235:1504-1508 (1987). The amino acid and DNA sequence of the M-CSF described therein is presented hereto in Fig. 1. Other forms of M-CSF bearing the active site thereof may also be employed in this composition, including synthetically produced polypeptides or polypeptides modified by recombinant means.

The term "M-CSF" is herein defined as including the naturally occurring human polypeptide M-CSF and naturally-

occurring allelic variations of the polypeptide. Allelic variations are naturally-occurring base changes in the species population which may or may not result in an amino acid change in a polypeptide or protein. Additionally included in this definition are both recombinant and synthetic versions of the polypeptide M-CSF, which may contain induced modifications in the peptide and DNA sequences thereof.

For example, the M-CSF polypeptide in the composition of the present invention may be characterized by a peptide sequence the same as or substantially homologous to the amino acid sequence illustrated in Fig. 1. These sequences may be encoded by the DNA sequence depicted in Fig. 1 or sequences containing allelic variations in base or amino acid sequence or deliberately modified structures coding for polypeptides with M-CSF biological properties.

Synthetic M-CSF proteins for use in the composition of the present invention may wholly or partially duplicate continuous sequences of the amino acid residues of Fig. 1. These sequences, by virtue of sharing structural and conformational characteristics with M-CSF polypeptides, e.g., the active site of the polypeptide of Fig. 1, may also possess M-CSF biological properties. Thus synthetic or recombinant polypeptides or fragments thereof may also be employed as biological or immunological equivalents for M-CSF polypeptides in the composition and methods of the present invention.

M-CSF, as used in the present invention also includes factors encoded by sequences similar to Fig. 1, but into which modifications are naturally provided or deliberately engineered. Modifications in the peptide or sequence of M-CSF can be made by one skilled in the art using known techniques. Specific modifications of interest in the M-CSF related sequences may include the replacement of one or more of the nine cysteine residues in the coding sequence with

other amino acids. Preferably several cysteines in each sequence are replaced with another amino acid, e.g. serine, to eliminate the disulfide bridges at those points in the protein. For example, lysine at amino acid position 163 (Fig. 1) could be deleted or substituted with another amino acid in order to eliminate the sensitivity of this region of M-CSF to trypsin-like proteases. Mutagenic techniques for such replacement are well known to one skilled in the art. [See, e.g., United States patent 4,518,584.]

Other specific mutations of the sequence of M-CSF described herein involve modifications of one or more of the glycosylation sites in the sequence. The absence of glycosylation or only partial glycosylation results from amino acid substitution or deletion at one, two, three or all of the asparagine-linked glycosylation recognition sites present in the sequence of M-CSF. The asparagine-linked glycosylation recognition sites comprise tripeptide sequences which are specifically recognized by appropriate cellular glycosylation enzymes. These tripeptide sequences are either asparagine-X-threonine or asparagine-X-serine, where X is usually any amino acid. A variety of amino acid substitutions or deletions at one or both of the first or third amino acid positions of a glycosylation recognition site (and/or amino acid deletion at the second position) results in non-glycosylation at the modified tripeptide sequence. Modification and variation of the types of oligosaccharides which attach to the O or N-linked glycosylation sites can occur by production of the sequence in either mammalian, bacterial, yeast or insect cells. Such modifications in the proteins are also encompassed by the term M-CSF.

Yet further modifications of M-CSF polypeptides may employ sequences which are designed for improved pharmacokinetics, by, e.g., association with polyethylene glycol. Alternatively, the last 25 to 35 amino acids of the

mature protein can be eliminated by appropriate gene deletion techniques to provide another form of M-CSF for use in the present invention. Such a deleted M-CSF may have use in genetic fusion to a cytotoxic agent. Amino acid residues 464 to 485 comprise a potential hydrophobic membrane-penetrating region. An M-CSF molecule that contains this sequence may desirably be employed in the composition of the invention, because these residues may embed the conjugate in the cell membrane, thereby aiding in the transfer of the cytotoxic agent into the cytosol.

An exemplary DNA sequence for the production of various M-CSF peptides have been deposited with the American Type Culture Collection, 12301 Parklawn Drive, Rockville, MD. The cDNA sequence illustrated in Fig. 1 below in vector p3ACSF-69, included in *E. coli* HB101 has been deposited on April 16, 1986 and given accession number ATCC 67092. This deposit was made under the Budapest Treaty on the International Recognition of the Deposit of Microorganisms for the Purposes of Patent Procedure and the Regulations thereunder (Budapest Treaty).

The cytotoxic agent linked to the M-CSF polypeptide is preferably a toxin or chemical agent which is capable of acting in the cytoplasm. Toxins may be employed which have a translocation property to move it through the cell membrane and a cytolytic domain, which provides its killing ability. One preferable class of toxins well-suited for this composition consists of two functionally different parts, termed A and B, which are connected by a disulfide bond. The A chain portion contains the enzymatic activity that enters the cytosol and kills the cell. The B chain moiety is responsible for binding of the toxin to the cell and presumably contains a domain that aids the A chain in crossing the cell membrane. Exemplary toxins for such use include native or genetically engineered ricin, abrin, modeccin, viscumin, Pseudomonas aeruginosa exotoxin,

Diphtheria toxin, Cholera toxin, Shigella toxin and E. coli heat labile toxin. The toxin portion of a conjugate prepared according to the invention can consist of the cytotoxic A chain portion only, the native holotoxin, or an engineered holotoxin, i.e., a toxin lacking its lectin binding property.

Other toxins which have only a single chain (an A chain portion) may also be employed. Examples of these toxins are ribosome inactivating proteins, such as pokeweed antiviral protein and gelonin. See, L. Barbieri et al, Cancer Surveys, 1:489-520 (1982) for a more complete list of ribosome inactivating proteins.

Mutant toxins or genetically engineered toxins may also be employed. Additionally microbially produced cytotoxic agents, and other non-protein organic molecules may be used as cytotoxic agents. The M-CSF ligand can also be linked to cytotoxic drugs, such as anthracyclines, e.g., doxorubicin, daunomycin, and the vinca alkaloids, such as, vindesine, vinblastine, vincristine. Methotrexate and its derivatives may also be employed as cytotoxic agents. More effective agents are those in which many molecules (between 5 to 50) of the drug are linked to the M-CSF through a polymer carrier, e.g., dextran. Bonds linking the drug to the carrier should be cleavable by the chemical environment inside the cell.

The M-CSF and a cytotoxic agent may be linked in a variety of ways. One way of linking these components is by employing one or more standard bifunctional protein crosslinkers, such as succinimidyl 3-(2-pyridyldithio)propionate (SPDP) or succinimidyl acetyl-thiopropionate (SATP). These crosslinkers form stable disulfide bonds between the M-CSF and toxin, or other cytotoxic agent, and yet are capable of releasing the toxin portion of the composition inside the cell, due to cleavage of the disulfide bonds by chemicals inside the cell, e.g., intracellular glutathione. These linking methods are known to those skilled in the art. See, e.g., J. Carlsson et al,

Biochem. J., 173:723-737 (1978) and N. Fujii et al, Chem. Pharm. Bull., 33:362-367 (1985). See also, A. J. Cumber et al, Methods Enzymol., 112:207-225 (1985) for other general methods for conjugating toxins to proteins.

For example, one method according to the invention involves making a M-CSF-toxin composition, using a toxin having both and A and B chain. The method involves the steps of:

(a) reacting the M-CSF with sufficient crosslinker to introduce between 1 to 6 reactive groups per molecule of M-CSF. A sufficient amount of crosslinker which can be used for this purpose is between approximately 6 to 50 moles of crosslinker per mole of M-CSF dimer.

(b) reacting a toxic protein having A and B chain subunits connected by at least one disulfide bond with a conventional reducing agent, thereby liberating the chains from each other.

(c) reacting the derivitized M-CSF of step (a) with the liberated A chain subunit of the reduced toxin; and

(d) separating from the reaction mixture conjugates comprising M-CSF linked by disulfide bonds to A chain subunits.

One exemplary growth factor/toxin conjugate is prepared by this method, modifying M-CSF with SPDP, followed by conjugation of ricin A chain toxin via a disulfide bond.

Another method for making the compositions of the present invention involves the following steps:

(a) reacting the M-CSF with sufficient crosslinker to introduce between 1 to 6 reactive groups per molecule of M-CSF;

(b) reacting the derivatized M-CSF of step (a) with a holotoxin having A and B subunits attached by at least one disulfide bond, the holotoxin being functionalized with a protein crosslinker which is preferably attached to the B subunit; and

(c) separating a conjugate formed by M-CSF becoming attached to the B subunit from free M-CSF and toxin in the reaction mixture.

Another manner of linking the components of the composition of the present invention is by a genetic fusion method. See, for example, United States Patent 4,675,382.

The compositions of the present invention containing both M-CSF and a toxin can be employed in methods for treating cancers characterized by over-expression of the c-fms proto-oncogene/M-CSF receptor gene. Among such cancers are acute myelocytic leukemia, ovarian cancer, breast cancer, lung cancer, pancreatic cancer and renal cancer. The composition of the invention operates by the targeting of the c-fms proto-oncogene by the M-CSF portion of the composition. Once attached to this receptor, the M-CSF molecule aids in transporting the cytotoxic agent through the cell membrane and into the cytosol. Inside the cell, the bonds linking the cytotoxic agent to the M-CSF are cleaved by chemicals naturally within the cell and the agent is released to kill the cancer cell.

The composition of the present invention can be administered in a variety of ways including systemically, locally or regionally. Desirably the composition is administered regionally in vivo, to the site of the carcinoma. For example, it can be administered intraperitoneally, if desired, to contain its distribution to the peritoneum for use in treating a suitable cancer, e.g., ovarian cancers. Similarly for treating lung cancers, the composition could be delivered in the form of an inhalant. If desirable, the composition may be administered subcutaneously, such as bathing effected tissue after surgical removal of a tumor e.g., for breast cancers. The composition may preferably be administered intravesically for instance into the bladder. Additionally, the composition can be employed in ex vivo applications, such as "purging" of a

mixture of cells removed from a patient, for patients having a systemic cancer which is not appropriate for regional application. The treatment of patients with acute myelocytic leukemia, for example, could involve removal of bone marrow cells from the body. These cells are then treated outside the body with the composition of the present invention to destroy a subset of these cells which are overexpressing the c-fms proto-oncogene. The "purged" cells are then reintroduced into the patient. The M-CSF/toxin composition of the invention can thereby serve as a purging agent to destroy the leukemic cells in the bone marrow of AML patients about to undergo autologous bone marrow transplantation. Other ex vivo purging treatments may also employ the composition of the invention.

The therapeutic composition for use in this invention may be in the form of a pyrogen-free, parenterally acceptable aqueous solution. The preparation of such a parenterally acceptable protein solution, having due regard to pH, isotonicity, stability and the like, is within the skill of the art.

The dosage regimen involved in treating the patient with the composition according to this invention will be determined by the attending physician considering various factors which modify the action of drugs, e.g. the condition, body weight, sex and diet of the patient, the severity of any infection, time of administration and other clinical factors. Additionally, the mode of administration could effect the dosage, e.g., ex vivo or in vivo. Generally, the daily regimen should be in the range of 2 to 2000 micrograms of polypeptide per kilogram of body weight.

The following examples illustrate the production of the M-CSF polypeptide and the construction of an M-CSF/toxin conjugate of the present invention.

EXAMPLE 1

Recombinant Production of M-CSF

To express the recombinant M-CSF polypeptide by recombinant means, the DNA encoding the polypeptide is transferred into an appropriate expression vector and introduced into selected host cells by conventional genetic engineering techniques.

Mammalian cell expression vectors for production of M-CSF, such as p3ACSF-69, may be synthesized by techniques well known to those skilled in this art. The components of the vectors, e.g. replicons, selection genes, enhancers, promoters, and the like, may be obtained from natural sources or synthesized by known procedures. See, Kaufman et al, J. Mol. Biol., 159:511-521 (1982); and Kaufman, Proc. Natl. Acad. Sci., U.S.A., 82:689-693 (1985). Suitable cells or cell lines for the expression of these recombinant M-CSF proteins may be Chinese hamster ovary cells (CHO), monkey COS-1 cells or CV-1 cells. The selection of suitable mammalian host cells and methods for transformation, culture, amplification, screening and product production and purification are known in the art. See, e.g., Gething and Sambrook, Nature, 293:620-625 (1981), or alternatively, Kaufman et al, Mol. Cell. Biol., 5(7):1750-1759 (1985) or Howley et al, U.S. Patent 4,419,446. Other exemplary mammalian host cells include particularly primate cell lines and rodent cell lines, including transformed cell lines. Normal diploid cells, cell strains derived from in vitro culture of primary tissue, as well as primary explants, are also suitable. Candidate cells need not be genotypically deficient in the selection gene so long as the selection gene is dominantly acting. For stable integration of the vector DNA, and for subsequent amplification of the integrated vector DNA, both by conventional methods, CHO cells may be employed. Other suitable mammalian cell lines include but are not limited to, HeLa, mouse L-929 cells, 3T3 lines

derived from Swiss, Balb-c or NIH mice, BHK or HaK hamster cell lines.

Stable transformants are then screened for expression of the product by standard immunological or enzymatic assays. The presence of the DNA encoding the variant proteins may be detected by standard procedures such as Southern blotting. Transient expression of the DNA encoding the variants during the several days after introduction of the expression vector DNA into suitable host cells such as COS-1 monkey cells is measured without selection by activity or immunologic assay of the proteins in the culture medium. The transformation of these vectors into appropriate host cells can result in expression of the M-CSF.

Similarly, one skilled in the art could manipulate the sequence of Fig. 1 by eliminating or replacing the mammalian regulatory sequences flanking the coding sequence with bacterial sequences to create bacterial vectors for intracellular or extracellular expression of M-CSF by bacterial cells. The DNA encoding the factor may be further modified to contain different codons for bacterial expression as is known in the art. Preferably the sequence is operatively linked in-frame to a nucleotide sequence encoding a secretory leader polypeptide permitting bacterial expression, secretion and processing of the mature variant protein, also as is known in the art. The compounds expressed in bacterial host cells may then be recovered, purified, and/or characterized with respect to physicochemical, biochemical and/or clinical parameters, all by known methods. For example, the M-CSF coding sequence could be inserted into a known bacterial vector using procedures such as described in T. Taniguchi et al., Proc. Natl Acad. Sci. USA, 77:5230-5233 (1980). This exemplary bacterial vector could then be transformed into bacterial host cells and the factor expressed thereby. The various strains of E. coli (e.g., HB101, MC1061) are well-known as

host cells in the field of biotechnology. Various strains of B. subtilis, Pseudomonas, other bacilli and the like may also be employed in this method. For a strategy for producing extracellular expression of such factors in bacterial cells, see, e.g. European patent application EPA 177,343.

Similar manipulations can be performed for the construction of an insect vector [See, e.g., procedures described in published European patent application 155,476] for expression in insect cells. See, e.g. Miller et al, Genetic Engineering, 8:277-298 (Plenum Press 1986) and references cited therein.

A yeast vector could also be constructed employing yeast regulatory sequences for intracellular or extracellular expression of M-CSF by yeast cells. Many strains of yeast cells known to those skilled in the art are also available as host cells for expression of the polypeptides useful in the invention. [See, e.g., procedures described in published PCT application WO 86 00639 and European patent application EP 123,289.]

EXAMPLE 2

An M-CSF Toxin Conjugate

For construction of an M-CSF toxin conjugate according to the invention, the growth factor M-CSF was produced in mammalian cells as described in pending U. S. patent SN940,362, the disclosures of which are incorporated by reference herein; and G. G. Wong et al, Science, 235 supra. M-CSF (5 mg, 55 nmoles) in 0.1M NaHCO₃ (20 ml) was reacted with a 20-fold molar excess of SPDP in ethanol. The reaction was allowed to proceed for five hours at 4 degrees Celsius to introduce approximately four to six sulfhydryl groups per molecule of M-CSF dimer. After removal of excess SPDP the derivatized growth factor was reacted with ricin A (15 mg, 500 nmoles), obtained from a commercial source, in 50mM NaH₂PO₄ p. 117.5/OIM NaCL. The disulfide bond was allowed

to form overnight at 4 degrees Celsius. The resulting M-CSF-ricin A chain conjugate was separated from excess ricin A chain by gel filtration on a Sepherogel™ TSK-3000 high pressure liquid chromatography column to give a mixture of conjugate and M-CSF (7.5 mg). After two passages through a column of Blue Sepharose developed with a gradient of NaCl, as described by P. P. Knowles and P. E. Thorpe, Anal. Biochem., 160: 440-443 (1987), the conjugate (720 mg) was obtained in a form free of M-CSF and consisted mainly of a species with one ricin A chain per M-CSF dimer.

EXAMPLE 3

In Vitro Cytotoxicity of M-CSF Toxin Conjugate

A level of toxicity and specificity for the M-CSF/ricin A chain conjugate was determined in a standard soft agar clonogenic assay in a manner similar to that described by Strong et al, Blood, 65: 627-635 (1985) with the NIH 3T3 and NIH 3T3-c-fms cell lines. The latter line which has been described by M. F. Roussel et al, Nature, 325: 549-552 (1987), is M-CSF receptor positive. Each cell line was mixed with either conjugate or medium without conjugate (control) in agarose and thin layered into Petri dishes. After incubation at 37°C in standard CO₂ atmosphere for a period of 14 days, the number of colonies in each dish was counted visually. The NIH 3T3-c-fms cells control dishes which did not receive the conjugate showed 103 colonies per dish while the same cells treated with conjugate at a concentration of 4 X 10⁻⁸M gave only 3 colonies. The NIH 3T3 cells, treated with conjugate and untreated control cells mixed with medium gave 76 and 78 colonies per dish, respectively.

EXAMPLE 4

Ex Vivo Assay of M-CSF Toxin Conjugate

The efficacy of the M-CSF/ricin A chain conjugate for ex vivo bone marrow purging is tested in a manner analogous to

that described by Strong et al, supra. M1 myeloid leukemic cells (10^3) which may be obtained from the American Type Culture Collection, Rockville, Maryland, (ATCC TIB 192) are added to murine bone marrow cells (10^5) and then treated with the M-CSF/ricin A conjugate in the 10^{-7} - $10^{-12}M$ range for approximately 4 hours at 37C. The percent survival of the leukemic cells as well as the monopotent and pluripotent bone marrow progenitor cells is determined with a standard colony formation assay, T.R. Bradley and D. Metcalf, Aust. J. Exp. Biol. Med. Sci., 44: 287 (1966) to measure the efficacy and specificity, respectively.

Numerous modifications may be made by one skilled in the art to the methods and components of the present invention in view of the disclosure herein. Such modifications are believed to be encompassed in the appended claims.

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International Application No: PCT/ /

MICROORGANISMS

Optional Sheet in connection with the microorganism referred to on page _____, line _____ of the description:

A. IDENTIFICATION OF DEPOSIT:Further deposits are identified on an additional sheet

Name of depository institution:

American Type Culture Collection

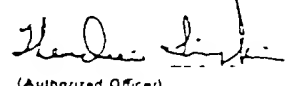
Address of depository institution (including postal code and country):

12301 Parklawn Drive
Rockville, Maryland 20852 USA

Name of Deposit	ATCC No.	Referred to on page/line	Date of Deposit
p3ACSF-69	67092	6/15	16 April 1986

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 (Authorized Officer)
 The date of receipt (from the applicant) by the International Bureau isWAS
(Authorized Officer)

WHAT IS CLAIMED IS:

1. A therapeutic composition for treating carcinoma characterized by over-expression of the c-fms proto-oncogene/M-CSF receptor gene comprising a M-CSF polypeptide conjugated to a cytotoxic agent and pharmaceutical carrier therefor.
2. The composition according to Claim 1, wherein said cytotoxic agent is a toxin selected from the group comprising double-chain ricin, ricin A chain, abrin, abrin A chain, modeccin and modeccin A chain, Pseudomonas aeruginosa exotoxin, Cholera toxin, Shigella toxin, E. coli heat labile toxin and Diphtheria toxin, mutant toxins thereof, and recombinant versions thereof.
3. The composition according to claim 1 wherein said cytotoxic agent is selected from the group consisting of ribosome-inactivating proteins, pokeweed antiviral protein and gelonin, mutant toxins thereof, and recombinant versions thereof.
4. The composition according to claim 1 wherein said cytotoxic agent is selected from the group consisting of anthracyclines, doxorubicin, daunomycin, vinca alkaloids, vindesine, vinblastine, vincristine, methotrexate and derivatives thereof.
5. The composition according to claim 1 where said M-CSF polypeptide is conjugated to said cytotoxic agent by a heterofunctional protein cross linking agent.
6. The composition according to claim 5 where said cross linking agent is selected from the group consisting of succinimidyl 3-(2-pyridyldithio)propionate) or succinimidyl

acetylthiopropionate.

7. The composition according to claim 1 comprising M-CSF conjugated through SPDP to a full ricin molecule.

8. A method for treating cancers characterized by an overexpression of the c-fms proto-oncogene/M-CSF receptor protein, comprising regionally administering in vivo to the site of said cancer a composition comprising M-CSF linked to a cytotoxic agent capable of crossing the membrane of the cell bearing said receptor and entering and killing the cell.

9. A method for treating cancers characterized by an overexpression of the c-fms proto-oncogene/M-CSF receptor protein, comprising ex vivo purging of a mixture of cells removed from a patient, said mixture containing said cancer cells, with a composition comprising M-CSF linked to a cytotoxic agent capable of crossing the membrane of the cell bearing said receptor and entering and killing the cancer cells.

10. A composition for treating carcinoma characterized by over-expression of c-fms proto-oncogene/M-CSF receptor gene comprising a monoclonal antibody to c-fms gene product/M-CSF receptor said monoclonal antibody conjugated to a cytotoxic agent and pharmaceutical carrier therefor.

Figure 1

10 20 30 40 50 60 70
 OCTGGGTCTT CTGGGCGCCA GAGGGCTCTT CGGCATCCCA GGACAGGGT GGGCGCTAG CGGGGGGGC

80 90 100 110 120 130 140
 CGACTCGGCA CGAGCGAGCG AGCGAGCGAG CGAGCGAGGG CGCGAGCGC CGCGCGCGG GACCGAGCTG

(-32) 160 175 190
 CGGTT ATG ACC GCG CGG GGC GGC GGG CGC TGC CCT CCC ACG ACA TGG CTG
 MET Thr Ala Pro Gly Ala Ala Gly Arg Cys Pro Pro Thr Thr Trp Leu

205 220 235 (1)
 GGC TCC CTG CTG TTG TTG GTC TGT CTC CTG GCG AGC AGG AGT ATC ACC GAG GAG
 Gly Ser Leu Leu Leu Leu Val Cys Leu Leu Ala Ser Arg Ser Ile Thr Glu Glu

250 265 280 295
 GTG TCG GAG TAC TGT AGC CAC ATG ATT CGG AGT GGA CAC CTG CAG TCT CTG CAG
 Val Ser Glu Tyr Cys Ser His MET Ile Gly Ser Gly His Leu Gln Ser Leu Gln

310 325 340 355
 CGG CTG ATT GAC AGT CAG ATG GAG ACC TCG TGC CAA ATT ACA TTT GAG TTT GTA
 Arg Leu Ile Asp Ser Gln MET Glu Thr Ser Cys Gln Ile Thr Phe Glu Phe Val

370 385 400
 GAC CAG GAA CAG TTG AAA GAT CCA GTG TGC TAC CTT AAG AAG CCA TTT CTC CTG
 Asp Gln Glu Gln Leu Lys Asp Pro Val Cys Tyr Leu Lys Lys Ala Phe Leu Leu

415 430 445 460
 GTA CAA GAC ATA ATG GAG GAC ACC ATG CGC TTC AGA GAT AAC ACC CCC AAT GCC
 Val Gln Asp Ile MET Glu Asp Thr MET Arg Phe Arg Asp Asn Thr Pro Asn Ala

475 490 505
 ATC GGC ATT GTG CAG CTG CAG GAA CTC TCT TTG AGG CTG AAG ACC TGC TTC ACC
 Ile Ala Ile Val Gln Leu Gln Glu Leu Ser Leu Arg Leu Lys Ser Cys Phe Thr

520 535 550 565
 AAG GAT TAT GAA GAG CAT GAC AAG GCG TGC GTC CGA ACT TTC TAT GAG ACA CCT
 Lys Asp Tyr Glu Glu His Asp Lys Ala Cys Val Arg Thr Phe Tyr Glu Thr Pro

580 595 (122) 610 625
 CTC CAG TTG CTG GAG AAG GTC AAG AAT GTC TTT AAT GAA ACA AAG AAT CTC CTT
 Leu Gln Leu Leu Glu Lys Val Lys Asn Val Phe Asn Glu Thr Lys Asn Leu Leu

640 655 670
 GAC AAG GAC TGG AAT ATT TTC AGC AAG AAC TGC AAC AAC AGC TTT GCT GAA TGC
 Asp Lys Asp Trp Asn Ile Phe Ser Lys Asn Cys Asn Asn Ser Phe Ala Glu Cys

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Figure 1 (Con't)

685 700 715 730
 TCC AGC CAA GAT GTG GTG ACC AAG CCT GAT TGC AAC TGC CTG TAC CCC AAA GGC
 Ser Ser Gln Asp Val Val Thr Lys Pro Asp Cys Asn Cys Leu Tyr Pro Lys Ala

745 760 775
 ATC CCT AGC AGT GAC CGC GCC TCT GTC TCC CCT CAT CAG CCC CTC GGC CCC TCC
 Ile Pro Ser Ser Asp Pro Ala Ser Val Ser Pro His Gln Pro Leu Ala Pro Ser

790 805 (189) 820 835
 ATG GGC CCT GTG GCT GGC TTG ACC TGG GAG GAC TCT GAG GGA ACT GAG GGC AGC
 MET Ala Pro Val Ala Gly Leu Thr Trp Glu Asp Ser Glu Gly Thr Glu Gly Ser

850 865 880 895
 TCC CTC TTG CCT GGT GAG CAG CCC CTG CAC ACA GTG GAT CCA GGC AGT GGC AAG
 Ser Leu Leu Pro Gly Glu Gln Pro Leu His Thr Val Asp Pro Gly Ser Ala Lys

910 925 940
 CAG CGG CCA CCC AGG AGC ACC TGC CAG AGC TTT GAG CGG CCA GAG ACC CCA GTT
 Gln Arg Pro Pro Arg Ser Thr Cys Gln Ser Phe Glu Pro Pro Glu Thr Pro Val

955 970 985 1000
 GTC AAG GAC AGC ACC ATC GGT GGC TCA CCA CAG CCT CGC CCC TCT GTC GGG GGC
 Val Lys Asp Ser Thr Ile Gly Ser Pro Gln Pro Arg Pro Ser Val Gly Ala

1015 1030 1045
 TTC AAC CCC CGG ATG GAG GAT ATT CTT GAC TCT GCA ATG GGC ACT AAT TGG GTC
 Phe Asn Pro Gly MET Glu Asp Ile Leu Asp Ser Ala MET Gly Thr Asn Trp Val

1060 1075 1090 1105
 CCA GAA GAA GCC TCT GGA GAG GGC AGT GAG ATT CCC GTC CCA CCC CAA CGG ACA GAG
 Pro Glu Glu Ala Ser Gly Glu Ala Ser Glu Ile Pro Val Pro Gln Gly Thr Glu

1120 1135 1150 1165
 CTT TCC CCC TCC AGG CCA GGA CGG CGC AGC ATG CAG ACA GAG CCC GGC AGA CCC
 Leu Ser Pro Ser Arg Pro Gly Gly Ser MET Gln Thr Glu Pro Ala Arg Pro

1180 1195 1210
 AGC AAC TTC CTC TCA GCA TCT CCT CCA CTC CCT GCA TCA GCA AAG GGC CAA CAG
 Ser Asn Phe Leu Ser Ala Ser Pro Leu Pro Ala Ser Ala Lys Gly Gln Gln

1225 1240 1255 1270
 CGG GCA GAT GTC ACT GGT ACA GGC TTG CCC AGG GTG GGC CCC GTG AGG CCC ACT
 Pro Ala Asp Val Thr Gly Thr Ala Leu Pro Arg Val Gly Pro Val Arg Pro Thr

1285 1300 1315
 GGC CAG GAC TGG AAT CAC ACC CCC CAG AAG ACA GAC CAT CCA TCT GGC CTG CTC
 Gly Gln Asp Trp Asn His Thr Pro Gln Lys Thr Asp His Pro Ser Ala Leu Leu

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Figure 1 (Con't)

1330 1345 1360 1375
 AGA GAC CCC CGG GAG CCA GGC TCT CCC AGG ATC TCA TCA CTG CGC CCC CAG GGC
 Arg Asp Pro Pro Glu Pro Gly Ser Pro Arg Ile Ser Ser Leu Arg Pro Gln Gly

1390 1405 1420 1435
 CTC AGC AAC CCC TCC ACC CTC TCT GCT CAG CCA CAG CTT TCC AGA AGC CAC TCC
 Leu Ser Asn Pro Ser Thr Leu Ser Ala Gln Pro Gln Leu Ser Arg Ser His Ser

1450 1465 1480
 TCG GGC AGC GTG CTG CCC CTT GGG GAG CTG GAG GGC AGG AGG AGC ACC ACG GAT
 Ser Gly Ser Val Leu Pro Leu Gly Glu Leu Gly Arg Arg Ser Thr Arg Asp

1495 1510 1525 1540
 CGG AGG AGC CCC GCA GAG CCA GAA GGA GGA CCA GCA AGT GAA CGG GCA CCC AGG
 Arg Arg Ser Pro Ala Glu Pro Glu Gly Pro Ala Ser Glu Gly Ala Ala Arg

1555 1570 1585
 CCC CTG CCC AGT TTT AAC TCC GTT CCT TTG ACT GAC ACA GGC CAT GAG AGG CAG
 Pro Leu Pro Arg Phe Asn Ser Val Pro Leu Thr Asp Thr Gly His Glu Arg Gln

1600 1615 1630 1645
 TCC GAG GGA TCC TCC AGC CGG CAG CTC CAG GAG TCT GTC TTC CAC CTG CTG GTG
 Ser Glu Gly Ser Ser Pro Gln Leu Gln Glu Ser Val Phe His Leu Leu Val

1660 1675 1690 1705
 CCC AGT GTC ATC CTG GTC TTG CTG GCT GTC GGA GGC CTC TTG TTC TAC AGG TGG
 Pro Ser Val Ile Leu Val Leu Ala Val Gly Leu Leu Phe Tyr Arg Trp

1720 1735 1750
 AGG CGG CGG AGC CAT CAA GAG CCT CAG AGA GCG GAT TCT CCC TTG GAG CAA CCA
 Arg Arg Arg Ser His Gln Glu Pro Gln Arg Ala Asp Ser Pro Leu Glu Gln Pro

1765 1780 1795 1817
 GAG GGC AGC CCC CTG ACT CAG GAT GAC AGA CAG GTG GAA CTG CCA GTG TAGAGGGAT
 Glu Gly Ser Pro Leu Thr Gln Asp Asp Arg Gln Val Glu Leu Pro Val

1827 1837 1847 1857 1867 1877 1887
 TCTTAAGCTGG ACGCACAGAA CAGTCTCTCC GTGGGAGGAG ACATTATGGG CGCTCCACCA CCACCCCTCC

1897 1907 1917 1927 1937 1947 1957
 CTGGCCATOC TCTCTGGAAATG TGGTCCTGCCC TCCACCAGAG CTCTGCTGTG CCAGGACTGG ACCAGAGCAG

1967 1977 1987 1997 2007 2017 2027
 CCAGGCCTGGG GCGCCCTCTGT CTCAACCGC AGAACCTTGA CTGAATGAGA GACGCCAGAG GATGCTCCCC

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Figure 1 (Con't)

2037 2047 2057 2067 2077 2087 2097
ATGCTGCCAC TATTTATTGT GAGCCTCGA GGCTCCCAATG TGCTTGAGGA AGGCCTGGTA GCGGGCTCA

2107 2117 2127 2137 2147 2157 2167
GGACOCTCTT CCTCAGGGG CTGCAACCTC CTCTCACTOC CTTOCATGCC GGAACCCAGG CCTAGGGACCC

2177 2187 2197 2207 2217 2227 2237
ACCGGCTGT CGTTTGIGGG AAAGCAGGGT GGACGCTGAG GAGTGAAGA ACOCTGCAOC CAGAGGCGCT

2247 2257 2267 2277 2287 2297 2307
GCCTGGTGOC AAGGTATOCG AGCCTGGACA GCCATGGACC TGTCCTCCAGA GAGAGGAGCC TGAAGTTGCT

2317 2327 2337 2347 2357 2367 2377
CCCCGGGAC AGGGTGGGCC TGATTTCGG TAAAGGTGIG CAGCTGAGA GAGGGAAAGA GGAGGCGCT

2387 2397 2407 2417 2427 2437 2447
GGACCTGGTG ACAGCTGAA GGGCTCACAC OCTGGCTCA OCTAAGTGOC CTGTCCTGGT

2457 2467 2477 2487 2497 2507 2517
TGCCAGGCGC AGAGGGGAGG CCTGGCTGAC CCTCAGGAOC TGCTGAOCCT GOCAGTGATG CCAAGAGGGG

2527 2537 2547 2557 2567 2577 2587
GATCAAGCAC TGGCTCTGC CCTCTCTCT TCCAGGACCT CCTAGAGCTT CCTCAGGAGG CCAAGCAGAG

2597 2607 2617 2627 2637 2647 2657
GCTCCCTCA TGAAGGAAGC CAITGCACTG TGAACACTGT ACOCTGCTGC TGAACAGGCT GCGGGGGTCC

2667 2677 2687 2697 2707 2717 2727
ATOCATGAGC CAGCATGGT CCTCTCTCA CTCTCCAGOC TCTCCCCAGC CTCTGCACT GAGCTGGCT

2737 2747 2757 2767 2777 2787 2797
CAOCACTGAGA CTGAGGGAGC CCTCAGGOC TGACCTCTC CTGACCTGGC CCTTGACTCC COGGAGTGGGA

2807 2817 2827 2837 2847 2857 2867
GTCGGGTGGG AGAAACCTCT GGGCGCGCAG CCTAGAGGAGG TCTTTAGGCT GTGTTGTTG OCAGGGTTTC

2877 2887 2897 2907 2917 2927 2937
TGCATCTTGC ACTTTGACAT TCCCAAGAGG GAAGGGACTA GTGGGAGAGA GCAAGGGAGG GGAGGGCACA

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Figure 1 (Con't)

2947 2957 2967 2977 2987 2997 3007
GACAGAGAGG AGCTCTGACT GAAGATGCC CTTTGAATA TAGGTATGCA CCTGAGGTTG

3017 3027 3037 3047 3057 3067 3077
GGGGAGGGTC TGCACTOOCA AACCCCCAGOG CAGTGTCCIT TOOCCTGCTGC CGACAGGAAC CTGGGGCTGA

3087 3097 3107 3117 3127 3137 3147
GCAGGTTATC CCTTGTCAGGA GCGCTGGACT GGGCTGCATC TCAGCCCCAC CTGCATGGTA TOCAGCTOOC

3157 3167 3177 3187 3197 3207 3217
ATCCACTTCT CAACCTTCIT TCTCTCTGAC CTTGGTCAGC AGTGATGAAC TCCAACTCTC ACCCCACCCCC

3227 3237 3247 3257 3267 3277 3287
TCTAACATCA CCTCTAAACCA CGCAAGOCAG CGTGGGAGAG CAATCAGGAG AGOCAGGCT CAGCTTOCAA

3297 3307 3317 3327 3337 3347 3357
TGCCCTGGAGG CGCTCAGCTT TGTTGGCAGC CTGTGGTGGT GGCTCTGAGG CCTAAGGCAAC GAGGAGCAGG

3367 3377 3387 3397 3407 3417 3427
GCTGOCAGTT CGCCCGCTGGT TCCCTTGTC TGCCTGTCGC CTCCTCTCT GCGCGCTTT GTCCTCGCT

3437 3447 3457 3467 3477 3487 3497
AAGAGACGCT CGCTAACCTG CGCGCTGGGC CGCGTGCCTT TCCCTCTG CGCAGGAAG TGAGGGTGG

3507 3517 3527 3537 3547 3557 3567
CTGGCCCCAC CTTCCTGTC CTGATGCGA CACCTTAGGG AAGGGCAGTG AACTTGCATA TGGGGCTTAG

3577 3587 3597 3607 3617 3627 3637
CCTTCTGACT ACAGCTCTA TATTTGATGC TAGAAAACAC ATATTTTAA ATGGAAGAAA AATAAAAAGG

3647 3657 3667 3677 3687 3697 3707
CATTCGGCTT TCATCCCCCT ACCCTAAACA TATAATATT TAAAGGTCAA AAAAGCAATC CAACCCACTG

3717 3727 3737 3747 3757 3767 3777
CAGAAGCTCT TTITGAGCAC TTGGTGGCAT CAGAGCAGGA GGAGCCCCAG AGOCACCTCT GGTGTCCCCC

3787 3797 3807 3817 3827 3837 3847
CAGGCTAACCT GCTCAGGAAC CGCTCTGTT CTCTGAGAAG TCAAGAGGG ACATGGCTC ACGCACTGTG

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Figure 1 (Con't)

3857 3867 3877 3887 3897 3907 3917
AGATTTTGT TTTATACCTG GAAGTGGIGA ATTATTTAT ATAAAGTCAT TAAATATCT ATTTAAAAGA

3927 3937 3947 3957 3967 3977
TAGGAAGCTG CTTATATATT TAATAATAAA AGAAGTGCAC AAGCTGCGT TGAGCTAGCT CGAG

INTERNATIONAL SEARCH REPORT

International Application No. PCT/US88/03697

I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) ⁶

According to International Patent Classification (IPC) or to both National Classification and IPC

Int. Cl 4 A61K 37/02, 31/705; C07K 17/06; C07H 15/24

U.S. CL 514/2, 8; 424/85.1; 530/351, 402, 403, 404, 405, 406

II. FIELDS SEARCHED

Minimum Documentation Searched ⁷

Classification System	Classification Symbols
U.S.	514/2, 8; 424/85.1; 530/351, 402, 403, 404, 405, 406

Documentation Searched other than Minimum Documentation
to the Extent that such Documents are Included in the Fields Searched ⁸

Computer Search on CAS and Dialog; Files CA, Biosis, 155,
350, 351; For: CSF and (conjugate or link or complex or
couple) and (toxin or cytotoxic agent or anthracycline)

III. DOCUMENTS CONSIDERED TO BE RELEVANT ⁹

Category ¹⁰	Citation of Document, ¹¹ with indication, where appropriate, of the relevant passages ¹²	Relevant to Claim No. ¹³
Y	US A, 4,504,586, (Nicolson), March 12, 1985, See Columns 1-2	1-8
Y	Science, Vol. 236, Issued June 1987; "The Human Hematopoietic Colony-Stimulating Factors", (Clark), pages 1229-37, See pages 1235-36.	1-8
Y	Blood, Vol. 67, Issued February 1986, "The Molecular Biology and Functions of the Granulocyte - Macrophage Colony - Stimulating Factors", (Metcalf), pages 257-67, See pages 259, 262-64.	1-8
Y	US, A, 4,675,382, (Murphy), June 23, 1987 See Columns 1-3.	1-2, 8-9
Y	Pharmac. Ther., Vol. 15, Issued 1982, "Chimeric Toxins", (Olsnes), pages 355-79, See pages 355, 357-62, 366.	1-8

* Special categories of cited documents: ¹⁰

"A" document defining the general state of the art which is not
considered to be of particular relevance

"E" earlier document but published on or after the international
filing date

"L" document which may throw doubt on priority claim(s) or
which is cited to establish the publication date of another
citation or other special reason (as specified)

"O" document referring to an oral disclosure, use, exhibition or
other means

"P" document published prior to the International filing date but
later than the priority date claimed

"T" later document published after the international filing date
or priority date and not in conflict with the application but
cited to understand the principle or theory underlying the
invention

"X" document of particular relevance; the claimed invention
cannot be considered novel or cannot be considered to
involve an inventive step

"Y" document of particular relevance; the claimed invention
cannot be considered to involve an inventive step when the
document is combined with one or more other such docu-
ments, such combination being obvious to a person skilled
in the art.

"&" document member of the same patent family

IV. CERTIFICATION

Date of the Actual Completion of the International Search

24 January 1989

Date of Mailing of this International Search Report

08 MAR 1989

International Searching Authority

Signature of Authorized Officer

Garnette D. Draper
Garnette D. Draper

III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)

Category *	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No.
Y	Exp. Clin. Cancer Res. Vol. 3, Issued 1984, "Biochemical Aspects of Antibody - Directed Delivery of Toxins and Drugs to Target Cancer Cells, (Chersi), pages 217-23.	1-8
Y	Monoclonal Antibodies '84: Biological and Clinical Applications, Issued 1985, "Antibody Carriers of Cytotoxic Agents in Cancer Therapy: A Review" (Thorpe), pages 475-506.	1-8